ATTORNEY DOCKET NO. 1568US/PCT

Pursuant to 37 CFR 1.8(a)(1)(i)(C), I hereby certify that this paper is being transmitted via the Office electronic filing system on <u>September 10</u>, 2008.

Carina Frazer

Docket No.: 1568US/PCT

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Patent No.: 7,202,470 Grant Date: April 10, 2007

Application No.: 09/787,358 371 Filing Date: 05/15/2001

Inventor: Philip Marriott PCT Filing Date: 09/16/1999

Title: MEANS FOR REMOVING UNWANTED IONS FROM AN ION TRANSPORT

SYSTEM AND MASS SPECTROMETER

Attention Certificate of Correction Commissioner for Patents P.O. Box 1450 Alexandria, VA 22313-1450

TRANSMITTAL

Transmitted herewith are the following documents:

Certification of Correction (1 pg.);

Copy of Excerpt from Response dated April 5, 2005

Pursuant to 37 C.F.R. §1.322 and MPEP §1480.01, Patentee requests expedited processing and granting of this request for a Certificate of Correction. In support of Patentee's assertion that the errors for which correction is sought are solely attributable to the Office, Patentee is submitting herewith a photocopy of the listing of claims submitted with the Response dated 04/05/05. It may be seen that claims 13, 14 (former claim 20), 28 (former claim 27), and 34 (former claim 67), as submitted, did not contain the errors for which correction is sought.

It is believed that no fee is due, however the Commissioner is hereby authorized to charge any appropriate fees under 37 C.F.R. §§1.16, 1.17, and 1.21 that may be required by this paper, and to credit any overpayment, to Deposit Account No. 50-3267.

Dated: September 10, 2008

Thermo Fisher Scientific Inc. ATTN: IP Department 355 River Oaks Parkway San Jose, California 95134 Tel: (408) 965-6000 Fax: (408) 965-6010 Respectfully submitted,

Charles B. Katz Reg. No. 36,564 Approved for use introducing destriction.

U.S. Patent and Trademark office; U.S. DEPARTIMENT OF COMMERCE

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(Also Form 2014).1601.

UNITED STATES PATENT AND TRADEMARK OFFICE CERTIFICATE OF CORRECTION

Page	4		4
		of	

PATENT NO. : 7,202,470

APPLICATION NO.: 09/787,358

ISSUE DATE : April 10, 2007

INVENTOR(S) : Philip Marriott

It is certified that an error appears or errors appear in the above-identified patent and that said Letters Patent is hereby corrected as shown below:

Claim 13, line 6

replace "analyte mass to charge ratio an ion optical device and" with --analyte mass to charge ratio and--

Claim 13, line 14-17

replace "analyte mass to charge ratio to produce a mass spectrum of the ion beam such that both the ion optical device and the mass analyzer operate at the same mass to charge ratio;" with –analyte mass to charge ratio:

Claim 13 line 20

replace "mass analyzing the received io beam" with --mass analyzing the received ion beam--

Claim 14, line 1

replace "A method according to claim 12" with --A method according to claim 13--

Claim 28, line 16-19

replace "receive ion beam to produce a mass spectrum of the received ion beam such that both the ion optical device and the mass analyzer operate at the same mass-to-charge ratio as the ion optical device" with --received ion beam at the same mass-to-charge ratio as the ion optical device-

Claim 34, line 1

replace "A mass spectrometer according to clamim 28" with --A mass spectrometer according to claim 28--"

MAILING ADDRESS OF SENDER (Please do not use customer number below):

Thermo Fisher Scientific Inc.

355 River Oaks Parkway

San Jose, CA 95134

This collection of information is required by 37 CFR 1.322, 1,323, and 1.324. The information is required to obtain or retain a benefit by the public which is to fixed by the USPTO to process) an episiciano. Condificativity is governed by \$5 U.S. C. 12 and \$7 CFR 1.14. This collection is estimated to flow to complete, including gathering, preparing, and equivalent to complete including gathering, preparing, and equivalent completed application form to the USPTO. Time will vary depending upon the individual case. Any comments on the second of time you require to complete is form and/or suppleasines for reducing the burden, should be sent to the Child is form and/or suppleasines for reducing the burden, should be sent to the Child is sent and the Suppleasines for reducing the burden, should be sent to the Child is sent and the suppleasines for reducing the burden, should be sent to the Child is commented to the sent to the Child is completed application. The comment of the sent to the Child is completed and the sent to the Child is completed and the sent to the Child is completed application. The sent the Child is completed and the sent the sent the Child is completed and the sent the Child is completed and the sent th

Attorney's Docket No.: 14544-002001 / P61716/003

Applicant : Philip Marriott Serial No.: 09/787.358 : May 15, 2001 Filed Page

: 2 of 15

Amendments to the Claims:

This listing of claims replaces all prior versions and listings of claims in the application:

Listing of Claims:

- 1. (Currently Amended) A mass spectrometer comprising: means (1) for generating ions from a sample introduced into a plasma;
- a sampling aperture (2) for transmitting some of the ions into an evacuated expansion chamber (3) along a first axis (9) to form an ion beam;
- a second aperture (5) for transmitting some of the ion beam into a first evacuated chamber (6):
 - a first pump (7) for maintaining the first evacuated chamber (6) at high vacuum;
- a first ion optical device (17) located in the first evacuated chamber (6) for containing the ion beam wherein the first ion optical device (17) is a mass selective device;
 - a third aperture (19) for transmitting the ion beam into a second evacuated chamber (20):
- a second pump (21) for maintaining the second evacuated chamber (20) at a lower pressure than the first evacuated chamber (6);
- a collision cell (24) having an entrance aperture (27) and an exit aperture (28) and pressurized with a target gas (26), the collision cell (24) being disposed in the second evacuated chamber (20);
- a second ion optical device (25) located in the collision cell (24) for containing the ion beam:
- a fourth aperture (32) for transmitting the ion beam into a third evacuated chamber (33) containing mass-to-charge ratio analyzing means (37) disposed along a second axis (36), wherein the mass-to-charge analyzing means is configured to mass analyze the ion beam to produce a mass spectrum of the ion beam such that both the first ion optical device (17) and the mass-tocharge ratio analyzing means (37) operate at the same mass to charge ratio, so as substantially to minimize the formation in the collision cell of interfering ions having the said mass to charge ratio:
 - a third pump (39) for maintaining the third evacuated chamber (33) at lower pressure than

Attorney's Docket No.: 14544-002001 / P61716/003

Applicant: Philip Marriott Serial No.: 09/787,358 Filed: May 15, 2001 Page: 3 of 15

the second evacuated chamber (20).

 (Original) A mass spectrometer according to claim 1, wherein the first evacuated chamber (6) is maintained at a pressure of approximately 10⁻² to 10⁻⁴ mbar.

- (Previously Presented) A mass spectrometer according to claim 1, wherein the first evacuated chamber (6) is maintained at a pressure of approximately 1-2 x 10³ mbar.
- (Previously Presented) A mass spectrometer according to claim 1, including a
 gap of at least 2 cm between the third aperture (19) and the entrance aperture (27) of the collision
 cell (24).
- (Previously Presented) A mass spectrometer according to claim 1, wherein the distance between the ion source (1) and the entrance aperture (27) of the collision cell (24) is 90 to 200 mm.
- (Previously Presented) A mass spectrometer according to claim 1, wherein the mass-to-charge ratio analyzing means (37) includes a main mass filter which preferably is an RF quadrupole.

7. (Cancelled)

- (Previously Presented) A mass spectrometer according to claim 1, wherein the first ion optical device (17) is an RF quadrupole.
- $9. \qquad \hbox{(Previously Presented) A mass spectrometer according to claim 1, wherein the second ion optical device (25) is an RF quadrupole.}$
- (Previously Presented) A mass spectrometer according to claim 1, wherein the second ion optical device (25) is mass selective.

Serial No.: 09/787,358
Filed: May 15, 2001
Page: 4 of 15

11. (Previously Presented) A mass spectrometer according to claim 1, wherein the second axis (36) of the mass to charge ratio analyzing means (37) is offset from the first axis (9).

- 12. (Previously Presented) A mass spectrometer according to claim 1, wherein the first evacuated chamber (6) is divided into a first region (14) adjacent to the expansion chamber containing an extractor lens (8) driven at a negative potential, and a second region (15) adjacent to the collision cell (24) in which the ion optical device (17) is located, by a large diameter aperture (11) and the aperture is sealable by means of a flat plate (12) on an O-ring seal (13).
- 13. (Currently Amended) A method of operating a mass spectrometer that incorporates a collision cell pressurized with a target gas, the method comprising:

generating, from an ion source, an ion beam by introducing a sample into a plasma, the ion beam including analyte ions having an analyte mass to charge ratio and artifact-unwanted ions;

mass selecting at least a portion of the ion beam at enthe analyte mass to charge ratio; transmitting at least a portion of the mass selected ion beam into the collision cell, the mass selecting step being effective substantially to minimize the formation in the collision cell of interfering ions having the analyte mass to charge ratio;

receiving at least a portion of the ion beam from the collision cell at a mass analyzer; and
mass analyzing the received ion beam at the same analyte mass to charge ratio as in the
mass selecting step.

transmitting at least a portion of the ion beam from the collision cell to a mass-analyzer; and

mass analyzing at least a portion of the ion beam in the mass analyzer at the analyte mass to charge ratio-

14. (Previously Presented) A method according to claim 13, wherein the mass selecting is achieved by passing the ion beam through a first mass selective ion optical device.

Serial No.: 09/787,358 Filed: May 15, 2001 Page: 5 of 15

 (Previously Presented) A method according to claim 14, wherein the first mass selective ion optical device is located in a first evacuated chamber maintained at high vacuum.

- 16. (Previously Presented) A method according to claim 15, wherein the collision cell is located in a second evacuated chamber operated at lower pressure than the first evacuated chamber, the ion beam being contained in the second evacuated chamber by a second ion optical device.
- (Previously Presented) A method according to claim 15, wherein the first evacuated chamber is maintained at a pressure of approximately 10⁻² to 10⁻⁴ mbar.
- (Previously Presented) A method according to claim 15, wherein the first evacuated chamber is maintained at a pressure of approximately 1-2 x 10³ mbar.
- 19. (Previously Presented) A method according to claim 16, further comprising transmitting at least a portion of the ion beam from the ion source through a sampling aperture into an evacuated expansion chamber along a first axis, into the first evacuated chamber through a second aperture;

wherein transmitting at least a portion of the mass selected ion beam into the collision cell includes transmitting at least a portion of the ion beam into the second evacuated chamber through a third aperture, wherein a gap of at least 2 cm is maintained between the third aperture and an entrance aperture of the collision cell.

- (Currently Amended) A method according to claim 13, wherein a distance of 90 to 200 emmm is maintained between the ion source and an entrance aperture of the collision cell.
- (Previously Presented) A method according to claim 19, wherein the mass analyzer is located in a third evacuated chamber operated at lower pressure than the second evacuated chamber, the mass analyzer being disposed along a second axis.

Serial No.: 09/787,358 Filed: May 15, 2001

Page : 6 of 15

 (Previously Presented) A method according to claim 14, wherein the first mass selective ion optical device is an RF quadrupole.

- (Previously Presented) A method according to claim 16, wherein the second ion
 optical device is an RF quadrupole.
- (Previously Presented) A method according to claim 16, wherein the second ion
 optical device is mass selective.
- 25. (Previously Presented) A method according to claim 15, wherein the first evacuated chamber is divided into a first region adjacent to the expansion chamber containing an extractor lens driven at a negative potential, and a second region adjacent to the collision cell, by a large diameter aperture and the aperture is sealable by means of a flat plate on an O-ring seal.
- (Previously Presented) A method according to claim 21, wherein the second axis
 is offset from the first axis.
 - 27. (Currently Amended) A mass spectrometer comprising:

an <u>plasma</u> ion source for generating an-ions beam-from a sample-introduced into a plasma;

an ion optical device disposed to receive at least a portion of an ion beam generated by the ion source, the ion optical device being configured to mass select at least a portion of the ion beam generated by the ion source at a mass-to-charge ratio;

a collision cell disposed to receive at least a portion of a mass selected ion beam from the ion optical device, the ion optical device being configured substantially to minimize the formation in the collision cell of interfering ions having the said mass-to-charge ratio; and

a mass analyzer disposed to receive at least a portion of the mass selected ion beam from the collision cell, the mass analyzer being configured to mass analyze the received ion beam at the <u>same</u> mass-to-charge ratio <u>as the ion optical device</u>.

Serial No.: 09/787,358 Filed: May 15, 2001

Page : 7 of 15

28-61. (Cancelled)

62. (New) A mass spectrometer according to claim 27, wherein the ion optical device and the mass analyzer are configured to scan synchronously.

- 63. (New) A mass spectrometer according to claim 27, wherein the mass analyzer is configured to mass select the ion beam received from the collision cell at the mass-to-charge ratio.
- (New) A mass spectrometer according to claim 27, wherein the ion optical device comprises a first RF quadrupole.
- (New) A mass spectrometer according to claim 64, wherein the mass analyzer comprises a second RF quadrupole.
- 66. (New) A mass spectrometer according to claim 27, wherein the ion optical device is disposed in a first evacuated chamber, the collision cell is disposed in a second evacuated chamber, and the mass analyzer is disposed in a third evacuated chamber.
- 67. (New) A mass spectrometer according to claim 27, further comprising a second ion optical device located in the collision cell for containing the ion beam.
- (New) A method according to claim 13, wherein mass selecting and mass analyzing comprise scanning synchronously.
- 69. (New) A mass spectrometer according to claim 1, wherein the first ion optical device and the mass-to-charge analyzing means are configured to scan synchronously.